Extracellular matrix production and calcium carbonate precipitation by coral cells in vitro

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The evolution of multicellularity in animals required the production of extracellular matrices that serve to spatially organize cells according to function. In corals, three matrices are involved in spatial organization: (i) an organic ECM, which facilitates cell–cell and cell–substrate adhesion; (ii) a skeletal organic matrix (SOM), which facilitates controlled deposition of a calcium carbonate skeleton; and (iii) the calcium carbonate skeleton itself, which provides the structural support for the 3D organization of coral colonies. In this report, we examine the production of these three matrices by using an in vitro culturing system for coral cells. In this system, which significantly facilitates studies of coral cell physiology, we demonstrate in vitro excretion of ECM by primary (non-dividing) tissue cultures of both soft (Xenia elongata) and hard (Montipora digitata) corals. There are structural differences between the ECM produced by X. elongata cell cultures and that of M. digitata, and ascorbic acid, a critical cofactor for proline hydroxylation, significantly increased the production of collagen in the ECM of the latter species. We further demonstrate in vitro production of SOM and extracellular mineralized particles in cell cultures of M. digitata. Inductively coupled plasma mass spectrometry analysis of Sr/Ca ratios revealed the particles to be aragonite. De novo calcification was confirmed by following the incorporation of 45Ca into acid labile macromolecules. Our results demonstrate the ability of isolated, differentiated coral cells to undergo fundamental processes required for multicellular organization.

Results and Discussion

Cell Culture Characterization. Extracellular production of organic matrices and calcium carbonate particles was examined in primary, nondividing cell cultures of the soft coral Xenia elongata and the stony coral Montipora digitata. Cell cultures were comprised of ~95% epithelium-like ectoderm or endoderm cells (the latter with or without zooxanthellae) ranging in size from 5 to 20 μm in diameter (Fig. 1), ~1% nematocysts, ~1% amoebocytes and <1% sensory nerve cells [supporting information (SI) Fig. 5]. The identification of cells was based on typical morphology and fluorescence (3). The cnidarian origin of cells in culture was verified by using 18S universal eukaryotic primers, and blasting the PCR-derived sequences against the National Center for Biotechnology Information nucleotide database (www.ncbi.nlm.nih.gov) (SI Fig. 6).

M. digitata cells were more granular, smaller, and less spherical than X. elongata cells (Fig. 1 b and a, respectively). The fraction of zooxanthellate/zooxanthellae-containing endoderm cells was always higher in the X. elongata cell cultures. However, after 2 weeks, chlorophyll fluorescence was virtually undetectable in both cell cultures. The suppression of chlorophyll fluorescence depended on the concentration of glucose in the medium; cells cultured at low glucose concentrations (0.1 mM glucose; Fig. 7) maintained high chlorophyll levels. These results suggest that glucose can suppress chlorophyll synthesis in zooxanthellae, possibly as a result of end-product inhibition (14–16). The viability of both M. digitata and X. elongata cell cultures remained >80% over a period of ~22 days (day 22: M. digitata, 83 ± 3%; X. elongata, 95 ± 3%).


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X. elongata, 87 ± 2%), after which it started to decrease (day 26: M. digitata, 70 ± 7%; X. elongata, 75 ± 6%). After the first week, many of the epithelial cells formed aggregates that adhered to the plates. Adhesion was best achieved on Primaria culture plates, which are coated with protonated amine groups and thus assume a positive charge. Because animal cells are characterized by a negative surface charge (17, 18), the initial attachment of cells to the plates presumably was attributable to electrostatic interactions.

**ECM Production.** The attached cells excreted ECM that mediated cell–cell as well as cell–substratum (plate surface) adhesion (Fig. 1 c–f). The ECM of both invertebrates and vertebrates is composed mainly of collagens, proteoglycans, and adhesive glycoproteins. The collagens (the fibrous structural proteins) are embedded in gels formed of polysaccharides (19, 20). SEM images of M. digitata and X. elongata cell aggregates indicate that the ECM formed in vitro indeed contained gel-like and fibrillar matrices (Fig. 1 c–f)

To examine the macromolecular composition of the ECM, we stained the cell cultures with Sirius red for collagen (21) and FITC–lectin conjugates of Con A and wheat germ agglutinin (WGA) for mannos/glucose and glucosamine/sialic acid residues in polysaccharides, respectively (22). These two classes of polysaccharides comprise the gel-like matrix in the ECM of _Pisaster ochraceus_ (phylum, Echinodermata) (22) and were shown to comprise the heteropolysaccharide in collagen from the sea anemone _Metridium dianthus_ (23). Sirius red, as well as Con A and WGA lectins, positively stained the ECM of both coral cell cultures (SI Fig. 8). Quantitative analysis (21) revealed that collagen accounted for 7.1 ± 0.7% of the total cell protein in _X. elongata_ cell cultures and 6.2 ± 0.5% of that in _M. digitata_.

Although WGA (SI Fig. 8) and Con A (data not shown) positively stained ECM and coral cell membranes, fluorometric quantification of polysaccharides in the ECM was not technically feasible.

Collagen is composed mainly of repeating glycine tripeptides with the sequence Gly-X-Y, where Y is frequently hydroxyproline (24). The hydroxylation of proline involves the activation of a reactive iron–oxygen complex, and ascorbic acid is required for the reduction of Fe$^{3+}$ formed in this reaction (25). Previous studies have shown that the addition of ascorbic acid stimulates collagen production in many metazoans (26–32); however, its effect on collagen production in basal metazoans such as corals is not known. The addition of ascorbic acid to the culture media increased collagen production by ~40% in _M. digitata_ cell cultures (4.4 ± 0.2 weight percent collagen relative to total protein without ascorbic acid compared with 6.2 ± 0.5% with ascorbic acid; P ≤ 0.01; Fig. 2); however, it had no significant effect on collagen production in _X. elongata_ cell cultures (6.0 ± 0.4% without ascorbic acid and 7.1 ± 0.7% with ascorbic acid; P ≥ 0.05). These results suggest either that collagen production in _X. elongata_ cell cultures, in contrast to _M. digitata_, is independent of ascorbate (33) or that ascorbate is not a limiting substrate for the formation of hydroxyproline. Indeed, when cells were cultured in the absence of ascorbic acid, the fraction of ECM–collagen of _X. elongata_ was significantly higher than that of _M. digitata_ cell cultures (P ≤ 0.05; Fig. 2).

**Calcium Carbonate Precipitation.** After ~2 weeks in culture, calcium carbonate-like particles, ranging in size from 20 to 100 µm, were visible in _M. digitata_ cell cultures (Fig. 3 and SI Fig. 9). Particles were observed within both adherent and nonadherent cell aggregates. The coverage of the particles by cells was not uniform and differed among the different aggregates (Fig. 3 and SI Fig. 9). Small mineral granules were also occasionally observed within _X. elongata_ cell aggregates; however, these granules, which were spherical in shape and ranged in size from 10 to 20 µm, were a relatively rare component and therefore were not analyzed further. The particles observed in _M. digitata_ cell cultures were identified as calcium carbonate based on the dominance of the Ca signal over other elements. High-resolution inductively coupled plasma mass spectrometry confirmed these
particles were aragonite based on Sr/Ca ratios (7.83 mmol mol\(^{-1}\); \(n = 10\)). Although this value is somewhat lower than that found in natural aragonitic coral skeletons (≈9.0 to 9.5 mmol mol\(^{-1}\)) (34), it is much higher than that of biogenic calcite (e.g., 1.15 to 1.45 mmol mol\(^{-1}\) for foraminifera) (35). We therefore conclude that the particles associated with the cell aggregates were a low-Sr form of aragonite.

To confirm that the aragonite was formed de novo, we measured incorporation of \(^{45}\text{Ca}\) into acid labile macromolecules. The calcification rates in \(M.\ digitata\) cell cultures were approximately an order of magnitude higher than those in \(X.\ elongata\) [1.8 ± 0.4 and 0.2 ± 0.08 nmol of Ca per milligram of protein per hour, respectively (mean ± SEM; \(P < 0.001\)]).

**Skeletal Organic Matrix.** Studies of coral biomineralization indicate that calcification is mediated by the synthesis of an organic framework that induces nucleation and crystal growth (4, 5, 36, 37). Analyses of the organic materials extracted from coral skeletons show a composition of acidic amino acids (mainly glutamic and aspartic acid) and sulfated polysaccharides (5, 38–41). To examine whether such an organic matrix exists within the calcium carbonate particles produced in vitro, cell aggregates were suspended in HCl for mineral digestion. Fig. 4 a–c shows a time series of particle dissolution by HCl, revealing a mucus-like organic matrix within the particles. Alcian blue is commonly used to detect mucopolysaccharides and glycoproteins (42) and was shown to stain sulfated polysaccharides and protein compounds extracted from coral skeletons (5). Staining the organic matrix within the particles with Alcian blue at pH 2.5 verified the polysaccharidic nature of the organic substance (Fig. 4d).

Despite progress in the field of coral biomineralization, information regarding the synthesis of the SOM and the pathway of calcification is still scarce. By examining the effect of various culture media on the composition of the SOM, we can better understand the processes affecting this pathway. Furthermore, although precipitation of aragonite is an extracellular process, the complexity of the coral skeleton prevents easy access to the interface compartments where calcification occurs, making in vivo studies of this process difficult. However, the relatively simple structure of the cell aggregates overcomes this problem by providing easy access to the extracellular location.

In vitro aragonite crystallization has been described previously in coral cell cultures of \textit{Pocillopora damicornis} (11). However, ECM and SOM production was not reported, and it was later shown that \textit{P. damicornis} exhibits an 80% decrease in cell viability after 7 days in culture (12). Dispersed single cells also failed to reaggregate or attach to the culture surface, suggesting that cell adhesion mechanisms were inactive (12). The medium used in this study is unique in that it is supplemented with aspartic and glutamic acids, which are major components of the coral SOM (40). Glutamine was supplemented as GlutaMAX (Invitrogen) a glutamine dipeptide that does not break down into ammonium, which can be toxic to cells (43). Furthermore, the constant addition of ascorbic acid to the culture medium may also aid in maintaining high cell viability because of its role as an antioxidant (44), as well as its importance for collagen production (24).

The results presented in this report reveal that coral cells maintain the ability to precipitate calcium carbonate on an ECM in vitro, while further excreting a matrix for cell–cell and cell–substratum organization. These processes are similar to bone formation and ECM production in higher metazoans. For example, both calcium carbonate precipitation in coral skeleton and calcium phosphate precipitation in bone result from mineral crystallization deposited on an organic matrix scaffold (45, 46). In addition, the SOM of both vertebrates and corals is composed mainly of acidic amino acids and polysaccharides (45, 46). The compatibility of coral skeleton as human implants (47, 48) further demonstrates the similarity between bone formation and coral calcification. Moreover, the ECM composition of corals is similar to that of higher metazoans (both in invertebrates as well as vertebrates) (19, 20, 49) and its production is considered to be a fundamental process in the evolution of multicellularity in animals (50). The cell culture system described here allows for detailed understanding of cellular processes that control calcification and ECM production in this basal metazoan group; the evolution of such processes were key events in the early evolution of metazoans.

**Materials and Methods**

**Corals.** Two coral species were used in this study: the soft coral \(X.\ elongata\) and the stony coral \(M.\ digitata\). Each species originated from one parent colony that was collected from reefs off of western Australia (Ashmore reef) and has been growing in an 800-liter, custom-designed aquarium as described previously (51).
Cell Cultures. To initiate cell cultures, small fragments of coral were excised from parent colonies and incubated for 2.5 h, with gentle shaking, in calcium-free artificial seawater supplemented with 3% antibiotics–antimycotics solution (GIBCO), prepared as described previously (12). The medium was replaced with artificial seawater 34% (Instant Ocean sea salt, Aquarium Systems) and 25 mM HEPES supplemented with collagenase (Sigma) at a final concentration of 1.5 mg/ml, and fragments were incubated for an additional 0.5 h. Fragments were then transferred to 35 × 10 mm Primaria culture dishes (two fragments in each dish; VWR International, Inc.). The plates contained 3 ml of the following culture medium: artificial seawater 34% (Instant Ocean sea salt, Aquarium Systems), 25 mM HEPES pH 8 – 8.0, 2% heat-inactivated FBS (Invitrogen), MEM vitamin solution 50×, MEM amino acid mix 100× (Sigma), GlutaMAX 2 mM (Invitrogen), aspartic acid 20 µg/ml, taurine 10 mM (Sigma), 1% antibiotic–antimycotics solution (GIBCO), 0.1–3 mM glucose, and 50 µg/ml L-ascorbic acid (added every other day). After 24 h, fragments were removed from the plates, and the medium with cells was passed (three times) through a custom-built sterile cell strainer. Cell strainers were made by cutting off a 15-ml Falcon tube into 3.5-cm-long columns and attaching, by heat, a 20-µm nylon mesh on one side. This insured the removal of tissue debris from the medium. Cells were maintained in a humidified chamber on a 12/12 h light/dark cycle at 26°C. After 24 h, cell strainers containing the coral fragments were removed from the plates. The medium was replaced with fresh medium, initially after 5 days and then every 7 days.

Microscopy Imaging. Light and fluorescence microscopy imaging was carried out with an inverted IX71 epifluorescence microscope (Olympus) equipped with a QImaging Retiga Exi SUVGA high-speed monochromatic cooled CCD camera system and IPLab for Mac (version 4.0.5) for image processing and analysis. For SEM of adherent cells, cells were fixed for 24 h with 2% formalin and gently washed with distilled water. After fixation, culture plates were cut into 1-cm2 pieces, followed by dehydration with an ascending ethanol series (70–100%) and critical point drying by using liquid CO2. Samples were then coated with gold and platinum and observed on an AMRAY-1830I microscope (Amaray, Inc.).

Cell Viability Measurements. Cell viability was quantitatively assessed by using Sytox green. Cells were incubated with 50 µM Sytox green for 15 min in the dark and visualized with an inverted IX71 epifluorescence microscope with FITC-/H9262 and GFP-/H11003 excitation and emission wavelengths. A total of 100–200 cells from three different optical fields of two culture plates were counted for each measurement (n = 6).

DNA Extraction 18S rDNA Amplification and Sequencing. For DNA extraction, cells were harvested and frozen in liquid nitrogen, followed by phenol/chloroform extraction and alcohol precipitation (52). 18S rDNA was amplified by PCR with general eukaryotic primers (5’-ACCGGTGTTTACCAATGCT-3’ and 5’-TGAATTCGACGCTCAG-3’) as described previously (53). PCR products were purified with the minielute gel extraction kit (Qiagen) and then cloned and transformed to TOP10 chemically competent bacteria by using the TOPO cloning kit (Invitrogen). Plasmids were purified by the QiAprep spin miniprep kit (Qiagen) and sequenced by using a 3100-Avant automatic sequencer (Applied Biosystems).

Quantification of Collagen and Total ECM Proteins. Collagen and proteins were quantified by colorimetric analyses by using Sirius red and Fast green as described previously (54). This method used the selective binding of Sirius red F3BA to collagen protein and Fast green FCF (Sigma) to noncollagen protein when both are dissolved in aqueous saturated picric acid. Briefly, culture plates were incubated with 1 ml of saturated picric acid solution that contained 0.1% Sirius red F3BA and 0.1% Fast green FCF. The plates were incubated at room temperature for 30 min in a rotary shaker. The fluids were then carefully withdrawn, and the plates were washed repeatedly with distilled water until the fluid was colorless. After washing, 1 ml of 11.1 (w/v) 0.1% NaOH and absolute methanol was added to the plates to elute the color. The eluted color was immediately read by using a spectrophotometer at 540 and 605 nm.

Lectin Staining. Cells were fixed with 2% formalin (w/v) before labeling with FITC–lectin conjugates of Con A and WGA (Sigma). After washing with artificial seawater, culture plates were incubated with 100 µl either lectin for 1.5 h at the dark. Plates were then washed twice to remove excess stain and were observed with an epifluorescence inverted microscope with FITC–suitable filters.

High-Resolution Inductively Coupled Plasma Mass Spectrometry. Precipitated mineral particles were pooled from three culture plates for inorganic analysis. The pooled cultures were incubated in 1 M NaOH at 90°C for 45 min to digest organic matter, then rinsed repeatedly with distilled water adjusted to pH 8 with NaOH. Particles were then rinsed with 0.065 M HNO3, the rinsing solution was removed by aspirating through a small pipette tip, and the particles were dissolved in 0.1 M HNO3 with sonication for 1 h (particles visibly dissolved). A 100-µl aliquot of this solution was diluted with 300 µl of 0.5 M HNO3 and analyzed for multiple elements, including Sr and Ca, by high-resolution inductively coupled plasma mass spectrometry against matrix-matched mixed-element standards (55).

Calcification Measurements. Calcification rates were measured as described in ref. 56 with modifications to adjust it for cell cultures (57). Briefly, cell cultures were incubated with 1 ml of culture medium containing 1 µCi of 45Ca (as CaCl2, 22.13 mM/ml; PerkinElmer) for 18 h. Cultures were maintained at 26°C under light (50 µmol of photons per meter squared per second) with gentle shaking (30 rpm). At the end of the incubation period, cells were scraped with a rubber policeman, and the medium containing calcium carbonate particles and cells was centrifuged at 20,000 × g for 2 min; the supernatant was retained for radioactive counting. Cells were washed until the supernatant did not contain radioisotope (five washes). Pellets were then resuspended in 0.5 ml of 1 M NaOH and incubated for 20 min at 90°C to dissolve organic tissue, followed by centrifugation at 20,000 × g for 2 min. The supernatant was kept for protein determination. The pellets were then washed with distilled water (pH 8) and resuspended with 0.5 ml of 6 M HCl for 18 h to dissolve the calcium carbonate particles. The samples were then added to 4.5 ml of scintillation liquid (Packard) and counted in a LS6000IC scintillation counter (Beckman).

Protein concentration was determined by using the BCA protein determination kit according to the protocol of the manufacturer (Pierce).

Exposure of Skeletal Organic Matrix and Alcian Blue Staining. Alcian blue solution was prepared in acetic acid 3% (10 g/liter). For decalcification, cultures were incubated in 1 M HCl until calcium carbonate particles had dissolved, followed by gentle washing with artificial seawater. Samples were then incubated with Alcian blue solution (pH 2.5) for 30 min, followed by washing with artificial seawater.

Statistical Analysis. Unless otherwise stated, all data were expressed as means ± SE. The Student’s t test was used to compare the differences between groups. A two-sample equal variance test was used when comparing the same coral species before and after treatment. A two-sample unequal variance test was used when comparing between the hard and the soft coral. Probability values of < 0.05 were considered significant.

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